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OM protein - protein search, using sw model

Run on: February 14, 2001, 03:34:30 ; Search time 36.01 seconds

(without alignments)
319.053 Million cell updates/sec

Title: US-09-481-990-2

Perfect score: 1753

Sequence: 1 MQLSLAGSSCVLVERHRSR.....QNEPVTAVQSSACVDPANH 336

Scoring table:

BLOSUM62

Gapop 10.0 , Gapext 0.5

Total number of hits satisfying chosen parameters: 268485

Minimum DB seq length: 0
Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%
Maximum Match 100%

Listing first 45 summaries

Database :

A.Geneseq_36:*

- 1: /cgn2_2/gcgdata/geneseq/geneseq/AA1980.DAT:*
- 2: /cgn2_2/gcgdata/geneseq/geneseq/AA1981.DAT:*
- 3: /cgn2_2/gcgdata/geneseq/geneseq/AA1982.DAT:*
- 4: /cgn2_2/gcgdata/geneseq/geneseq/AA1983.DAT:*
- 5: /cgn2_2/gcgdata/geneseq/geneseq/AA1984.DAT:*
- 6: /cgn2_2/gcgdata/geneseq/geneseq/AA1985.DAT:*
- 7: /cgn2_2/gcgdata/geneseq/geneseq/AA1986.DAT:*
- 8: /cgn2_2/gcgdata/geneseq/geneseq/AA1987.DAT:*
- 9: /cgn2_2/gcgdata/geneseq/geneseq/AA1988.DAT:*
- 10: /cgn2_2/gcgdata/geneseq/geneseq/AA1989.DAT:*
- 11: /cgn2_2/gcgdata/geneseq/geneseq/AA1990.DAT:*
- 12: /cgn2_2/gcgdata/geneseq/geneseq/AA1991.DAT:*
- 13: /cgn2_2/gcgdata/geneseq/geneseq/AA1992.DAT:*
- 14: /cgn2_2/gcgdata/geneseq/geneseq/AA1993.DAT:*
- 15: /cgn2_2/gcgdata/geneseq/geneseq/AA1994.DAT:*
- 16: /cgn2_2/gcgdata/geneseq/geneseq/AA1995.DAT:*
- 17: /cgn2_2/gcgdata/geneseq/geneseq/AA1996.DAT:*
- 18: /cgn2_2/gcgdata/geneseq/geneseq/AA1997.DAT:*
- 19: /cgn2_2/gcgdata/geneseq/geneseq/AA1998.DAT:*
- 20: /cgn2_2/gcgdata/geneseq/geneseq/AA1999.DAT:*
- 21: /cgn2_2/gcgdata/geneseq/geneseq/AA2000.DAT:*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

SUMMARIES

Result No.	Score	Query Match	Length	DB ID	Description
1	1753	100.0	336	18 W23397	TIWK-1 potassium c
2	653.5	37.3	313	20 Y34132	Human potassium ch
3	653.5	37.3	313	20 Y25116	Human hTREK-1 prot
4	653.5	37.3	313	21 Y68737	KT4, a TWIK family
5	653.5	37.3	313	21 Y68738	KT5, a TWIK family
6	379.5	21.6	398	20 Y30647	A mechanically sen
7	379.5	21.6	411	20 Y34133	Human potassium ch
8	379.5	21.6	411	20 Y28496	h-TREK1 polypeptid
9	378	21.6	393	21 Y94425	Human h-TRAAK poly
10	378	21.6	393	21 Y94426	Human h-TRAAK poly
11	376.5	21.5	370	20 Y30648	A mechanically sen
12	376.5	21.5	411	20 Y28497	Mouse h-TREK1 poly

13	359	20.5	499	21 Y94875
14	320.5	18.3	194	21 Y87291
15	315	18.0	197	20 Y34126
16	239	13.6	618	17 R97984
17	220	12.5	336	17 R97986
18	163	9.3	107	20 Y28498
19	118.5	6.8	417	20 Y32010
20	118.5	6.8	417	20 Y32010
21	116.5	6.6	962	21 Y49944
22	116.5	6.6	989	21 Y49945
23	113.5	6.5	1102	21 Y83028
24	111	6.3	1107	21 Y44907
25	110	6.3	597	20 Y32012
26	107	6.1	542	21 Y44905
27	98.5	5.6	843	20 Y06561
28	98.5	5.6	843	20 W99799
29	97.5	5.6	252	19 W74743
30	97.5	5.6	857	20 Y32019
31	95.5	5.4	434	17 R92315
32	95	5.4	858	20 Y32015
33	94.5	5.4	363	16 R66934
34	94	5.4	494	17 R90765
35	94	5.4	494	19 W42996
36	94	5.4	1017	20 Y22427
37	93.5	5.3	425	20 W98019
38	93.5	5.3	1174	11 R08257
39	93.5	5.3	1174	13 R25825
40	93.5	5.3	1174	17 R89494
41	93.5	5.3	1174	18 W09043
42	92	5.2	495	20 Y33766
43	91.5	5.2	461	18 W21009
44	91.5	5.2	1159	20 Y32020
45	91.5	5.2	1159	21 Y85405

ALIGNMENTS

RESULT 1	
W23397	W23397 standard; Protein; 336 AA.
XX	
AC	W23397;
XX	
DT	17-MAR-1998 (first entry)
XX	
DE	TIWK-1 potassium channel protein.
XX	
KW	TIWK-1 potassium channel; screening; diagnosis; transgenic animal;
KW	Tandem of P domains in a weak inward rectifying K ⁺ antibody.
XX	
OS	Homo sapiens.
XX	
FN	FR2744730-A1.
XX	
PD	14-AUG-1997.
XX	
XX	
PF	08-FEB-1996; 96FR-0001565.
XX	
PR	08-FEB-1996; 96FR-0001565.
XX	
PA	(CNRS) CNRS CENT NAT RECH SCI.
PI	Barhanin J, Duprat F, Fink M, Guillemaire E, Lazdunski M;
PI	Lesage F, Romey g;
XX	
DR	WPI: 1997-427773/40.
XX	
XX	N-PSDB; T64960.
PT	Nucleic acid encoding new potassium channel designated TWIK-1 -
PT	Human h-TRAAK poly
PT	active agents and for diagnosis

Human protein clon
Human signal pept
Human potassium ch
DMORF1 potassium c
F22b7.7 potassium
partial h-TREK1 po
Escherichia coli c
Rat Bag1 potassium
Human potassium io
Human potassium io
Rat Elki potassium
Human potassium ch
Paramecium tetrau
Human potassium ch
Chicken capsaicin
Chicken VRI capsa
Human secreted pro
Arabidopsis thalia
CORK potassium cha
Human cation chann
Mouse ATR2 receptor
Human K⁺ channel 2
Putative mature po
Human brain specif
Mouse calcium acti
B. thuringiensis to
Novel toxin expres
B. t. toxin 81A2.
Bacillus thuringie
hkv5.1 human brain
H. pylori cell env
Human cation chann
Long QT syndrome a

RESULT 3
 Y25116
 ID Y25116 standard; Protein; 313 AA.
 AC Y25116;
 DT 25-AUG-1999 (first entry)
 XX
 DE Human hTREK-1 protein.
 KW hTREK-2; Twik-1 Related K⁺ channel-2; vasotropic; antiinflammatory;
 KW analgesic; treatment; gene therapy; inhibitor; detection; diagnosis;
 KW disease susceptibility; cerebral; cardiac; renal; ischemia; brain;
 KW inflammation; pain; mimic; neurotransmitter; hormone; chromosome mapping
 KW linkage analysis; mutation; immunogen; human.
 XX
 OS Homo sapiens.
 PN EP930364-A1.
 XX
 PD 21-JUL-1999.
 XX
 PE 16-JAN-1998; 98EP-0400072.
 XX
 PR 16-JAN-1998; 98EP-0400072.
 XX
 PA (SYNO) SYNTHELABO.
 XX
 PI Partisetti M;
 DR WPI; 1999-387707/33.
 DR N-PSDB; X78383.
 XX
 PT New human polypeptides useful for diagnosing and treating cerebral
 PT and cardiac ischemias .
 XX
 PS Claim 1; Page 18-20; 21pp: English.
 XX
 AB This invention describes a novel human Twik-1 Related K⁺ channel-2
 (hTREK-2) polypeptide (I) and its encoding nucleic acid (II) which has
 vasotropic, antiinflammatory and analgesic activity. (II) or agonists of
 (I) may be used to stimulate production of (I) in vivo to treat patients
 requiring enhanced activity or expression of (I). This use of (II)
 represents a gene therapy regime. Antagonists of (I), the complement of
 (II) used as an antisense construct or a polypeptide competitor of (I)
 may be administered to patients to inhibit activity or expression of (I)
 detection of the presence or amount of (I) in a sample from a patient or
 detection of mutations in (I) may be used to diagnose or measure
 susceptibility to diseases related to altered expression or activity of
 (I). The diseases and conditions resulting from altered activity or
 expression of (I) which may be treated as above include cerebral,
 cardiac and renal ischemias, brain and cardiac diseases, inflammation
 and pain. In addition, (I), (II), and agonists and antagonists of (I)
 may be used to mimic or antagonize the effects of endogenous
 neurotransmitters and hormones. (II) or its fragments may be used as
 hybridization probes to isolate full length and genomic cDNAs encoding
 (I) or its homologues from cDNA or genomic libraries. (II) may also be
 used for chromosome mapping and linkage analysis to identify the
 relationship between genes and diseases which have been mapped to the
 same chromosome. In addition (II) may be used to identify mutations
 associated with diseases by comparing the sequence of (II) between
 affected and unaffected individuals. (I) or its fragments may be used as
 immunogens to produce antibodies against (I). Antibodies to (I) may be
 used to isolate or identify clones expressing (I) or to purify (I) by
 affinity chromatography. These antibodies may also be used to treat the
 above diseases as agonists or antagonists of (I).
 XX

Query Match	37.38;	Score 653.5;	DB 20;	Length 313;
Best Local Similarity	45.38;	Pred. No. 3.2e-60;		

Matches	140;	Conservative	58;	Mismatches	86;	Indels	25;	Gaps	7;
QY	24	GFVLVG----	YLLVYVSGAVVFSVELPYEDLLRQELKKRLRLEEHCCSEQOLEQFL	79					
Db	4	gallagalaayaaylvlgallvarleqphearleaeletlraqllqrpvcvaapaldafv	63						
QY	80	GRVLEASNVGVSVLSNAGSNMN-----WDFSALEPFASTVLSTGCGTHTVPLSDGKAFIC	134						
Db	64	ervlaaagrlgrvrvlaanaagsnaadqadafasalfifastllltvygytclptldgkafis	123						
QY	135	IIVSVIGIPFLLLEFATAVQRTVTVTRRPVLYEFHIRMGFSKOVAYIAHAV-LIGFVTVS	193						
Db	124	lafalagpctmllltasaagrlsltlthvpslwmrgwqpprraacchlyallagvvtv	183						
QY	194	CFEFPDPAVFSVLEDDWNFLSEFYFCFSTSLSTIGLDGVYPGEGYNQKRELYKIGITCYL	253						
Db	184	c-flvpavilafahleawsfldafyfcflstlctlgdyvpgpaqbpqpylvtvyl	242						
QY	254	LLGLTAMLVVETPELHELKKRFMFV-----KKDDEQVHII-----EHDLDS	300						
Db	243	flglvawmvlvqtrfhvsgdlngltelillppcpasfnaded-drvdlilqpseshqqls	301						
QY	301	FSSITPDQAA	309						
Db	302	asshtcdyas	310						
RESULT	4								
Y68737	ID	Y68737	standard; Protein; 313 AA.						
AC	Y68737;								
XX	05-MAY-2000	(first entry)							
DT									
XX									
DE	KT4, a TWIK family 2PD potassium channel polypeptide.								
XX									
KW	KT4, TWIK family 2PD potassium channel polypeptide; P-domain;								
KW	expressed sequence tag; EST: AA604914; ion channel dysfunction;								
KW	renal failure; musculoskeletal disease; proliferative disease;								
KW	renal failure; nephrosis; cirrhosis; dysphagia; gastritis; myoclonia;								
KW	muscular dystrophy; atherosclerosis; cancer.								
XX									
OS	Homo sapiens.								
XX									
PN	WO200003687-A2.								
XX									
PD	27-JAN-2000.								
XX									
PF	20-JUL-1999;	99WO-US16471.							
XX									
PR	20-JUL-1998;	98US-0093486.							
PR	13-AUG-1998;	98US-0096655.							
XX									
PA	(ELAN-) ELAN PHARM INC.								
XX									
PI	Forsayeth JR, Zhao BB, Chavez RA;								
XX									
DR	WPI: 2000-171196/15.								
XX	N-PSDB; 246092.								
PT									
PT	Novel human potassium channel polynucleotides and polypeptides used in								
PT	the diagnosis, prevention and treatment of diseases including renal								
PT	failure, cirrhosis, muscular dystrophy and cancers .								
XX									
PS	Claim 4; Fig 1A-C; 53pp: English.								
XX									
CC	The present sequence represents a protein, designated KT4, which is a								
CC	member of the TWIK family 2PD potassium channel polypeptides. These								
CC	polypeptides contain two potential P-domains and 8 (preferably 4)								
CC	transmembrane domains. The KT4 cDNA sequence was isolated from a brain								
CC	cDNA library using degenerate oligonucleotides derived from human								
CC	expressed sequence tag (EST) AA604914. The polypeptides and								

RESULT	4
ID	Y68737
XX	Y68737 standard; Protein; 313 AA.
XX	
AC	Y68737;
XX	
DT	05-MAY-2000 (first entry)
XX	
DE	KT4, a TWIK family 2PD potassium channel polypeptide.
XX	
KW	KT4, TWIK family 2PD potassium channel polypeptide; P-domain;
KW	expressed sequence tag; EST; AA604914; ion channel dysfunction;
KW	renal disease; musculoskeletal disease; proliferative disease;
KW	renal failure; nephrosis; cirrhosis; dysphagia; gastritis; myotonia;
KW	muscular dystrophy; atherosclerosis; cancer.
XX	
OS	Homo sapiens.
XX	
PN	WO200003687-A2.
XX	
PD	27-JAN-2000.
XX	
PF	20-JUL-1999; 99WO-US16471.
XX	
PR	20-JUL-1998; 98US-0093486.
PR	13-AUG-1998; 98US-009655.
XX	
PA	(ELAN-) ELAN PHARM INC.
XX	
PI	Forsayeth JR, Zhao BB, Chavez RA;
XX	
DR	WPI: 2000-171196/15.
XX	N-PSDB; Z46092.
XX	
PT	Novel human potassium channel polynucleotides and polypeptides used in
PT	the diagnosis, prevention and treatment of diseases including renal
PT	failure, cirrhosis, muscular dystrophy and cancers -
XX	
PS	Claim 4; Fig 1A-C; 53pp; English.
XX	
CC	The present sequence represents a protein, designated KT4, which is a
CC	member of the TWIK family 2PD potassium channel polypeptides. These
CC	polypeptides contain two potential P-domains and 8 (preferably 4)
CC	transmembrane domains. The KT4 cDNA sequence was isolated from a brain
CC	cDNA library using degenerate oligonucleotides derived from human
CC	expressed sequence tag (EST) AA604914. The polypeptides and
CC	

CC polynucleotides are used in the diagnosis, prevention and treatment of
CC disease states. The polynucleotides may be used to detect and
CC quantitative expression of TWIK family 2PD potassium channels, and
CC aberrant or mutant forms of the polynucleotide which cause various
CC diseases and disorders. Antisense oligonucleotides may be used to
CC modulate the expression of polynucleotides of the invention. The
CC polypeptides are used for treating diseases and disorders associated
CC with ion channel dysfunction, including renal, musculoskeletal and
CC proliferative diseases, e.g., renal failure, nephrosis, cirrhosis,
CC dysphagia, gastritis, myotonia, muscular dystrophy, atherosclerosis
CC and cancers.

50	Sequence	313	AA;
----	----------	-----	-----

Query Match	37.38;	Score 653.5;	DB 21;	Length 313;
Best Local Similarity	45.28;	Score 733.50;	DB 21;	Length 313;

Matches 140; Conservative 58; Mismatches 86; Indels 25; Gaps 7;

QY 24 GFLVLG---YLIVYGVGAFFSSVEPYEDLLKQELRKIKRPLENHECLSDQEQFL 79
4 gallagalaayaaglylgallvarleagphnearlraaelcragllqpscevaapadaty 63

QY 80 GRVLEASNTGYSVLSNAGSMWN----WDFTSALFPASTVLTSTTGXHTVPLSDGKAFIC 134
64 etvlaeaqrllgrvvlaanagsanaspawdfasallfcastllttvyguytcepltdagafas 1233

QY 135 IITSVIGIPPTLEFLFVAVVORITVHVRRPVLEPHNMGSKQVVALVNHV-LLGPTVS 1233
124 fatallgyvptcmlllttsaqrlslllthpvlswmvgwdprtraacwhllyallgvvv 1833

QY 194 CFFFTLPAAVSVLEDDMNFLESTFFTCISLSTIGLDGYDPGEGYNOKFRELYIGITCYL 2533
184 c-flvpevafiahleaaasfldayfclstlsglygdyvpgaeapqpyralykvlyl 242

QY 254 ILGLIALVYLEFCELEHKKFRKMYV-----KKRKDEOVYII-----EHDQLS 300
243 flglvamvlylgcfhrhsdnhgtellllpppocastnadcd-drydvlpgpreshqds 301

QY 301 FSSSTPDQA 309
302 assthtdyas 310

RESULT	5
Y68738	
ID	Y68738 standard; Protein; 313 AA

05-MAY-2000 (first entry)

KT5, a TWIK family 2PD potassium channel polypeptide.

renal disease: musculoskeletal disease, proliferative disease, renal failure, nephrosis, cirrhosis, cataracts, cancer, muscular dystrophy, atherosclerosis; dysc.

OS Homo sapiens.

PN WO200003687-A2.

PD 27-JAN-2000.

PF 20-JUL-1999; 99WO-US16471.

PR	20-JUL-1998;	98US-0093486.
PR	13-AUG-1998	00US-0005555

PK 13-AUG-1998; 9805-0096655.
XY

(ELAN-) ELAN PHARM INC.

PI Forsayeth JR, Zhao BB, Chavez RA;
v

DR WPI; 2000-171196/15.

DR N-PSDB; 246094.

XX failure, cirrhosis, muscular dystrophy and cancers -
 PT Novel human potassium channel polynucleotides and polypeptides used in
 PT the diagnosis, prevention and treatment of diseases including renal
 PS Claim 7; Fig 3A-C; 53pp; English.

The present sequence represents a protein, designated K15, which is a member of the TWIK family 2Pb potassium channel polypeptides. These polypeptides contain two potential P-domains and 8 (preferably 4) transmembrane domains. The K14 cDNA sequence was isolated from a Brain cDNA library using degenerate oligonucleotides derived from human expressed sequence tag (EST) A45312. The polypeptides and polynucleotides are used in the diagnosis, prevention and treatment of disease states. The polynucleotides may be used to detect and quantitate expression of TWIK family 2Pb potassium channels, and aberrant or mutant forms of the polynucleotide which cause various diseases and disorders. Antisense oligonucleotides may be used to modulate the expression of polynucleotides of the invention. The polypeptides are used for treating diseases and disorders associated with ion channel dysfunction, including renal, musculoskeletal and proliferative diseases, e.g., renal failure, nephrosis, cirrhosis, dysphagia, gastritis, myotonia, muscular dystrophy, atherosclerosis and cancers.

Sequence 313 AA;

Query Match	37.38;	Score 653.5;	DB 21;	Length 313;
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Matches	140;	Conservative	58;	Mismatches	86;	Indels	25;	Gaps	7;
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QY	24	GFVIVLG----	YLYLVVFVFAVSSVEVPEYDDLROELRKRRFLREHHCSEQLEOFL	79	
Db	4	galigalaaayayvvgallvarleoghpearlraelctrlqllqirspcvaapaladfv	63		
QY	80	GRVLEASNTGYSVLSNMSGMN----	WDTSLAFPASTVLTSTGYGHTVPLSDGGAFC	134	
Db	64	ervlaagrlgrvrvlanaasanaadpawdfasalafastlltvgygvtptldagkafs	123		
QY	135	IIVSYGIPFELLFTTAVVGRITVHWRRVVLFEHFRMGESKOVAIVAAV-LIGFVTVS	192		
Db	124	isfallgpytcmllltsaagllslllthpyslsmrwspopraacwhlvalllgyvvvtv	183		
QY	194	CEFFTPAAVFSVLEDDNNFLFESFPFCISLSTGGLDYYPGSGYQOKFRELYKIGITCYL	253		
Db	184	c-flvpavflahleavsfldafyfctislsistlgldgvpgaapqpyralykvltvyl	242		
QY	254	LGLTAMLVVLETCFCEHLKFRKMFV-----	KDKREDVNIIT-----	EHDLIS	300
Db	243	flgryamvlvqtrfnrshdngltellllppropasfaded-drvdlldgqpesbqqls	301		
QY	301	ESSITDOAA	309		
Db	302	ashtcdyaa	310		

RESULT	6
Y30647	
ID	Y30647 standard; Protein; 398 AA.

18-NOV-1999 (first entry)

A mechanically sensitive potassium channel protein TRAAK

KM Mechanically sensitive potassium channel protein; TRAAK;
 KM polynsaturated fatty acid; arachidonic acid; riluzole; heart disease

KW nervous system disease; epilepsy; cardiovascular disease; arrhythmia;
 KW neurodegeneration; ischemia; anoxia; hormone secretion abnormality;
 KW muscular disease.
 XX Mus sp.
 OS WO9945108-A2.
 XX PD 10-SEP-1999.
 XX PF 23-FEB-1999; 99WO-FR00404.
 XX PR 05-MAR-1998; 98FR-0002725.
 XX PA (CNRS) CNRS CENT NAT RECH SCI.
 XX PI Honore E, Fink M, Lazdunski M, Lesage F, Duprat F;
 WPI: 1999-551038/46.
 N-PSDB: 210606.
 XX PT New mechanically sensitive potassium channel, used to screen for
 PT specific modulators, potential therapeutic agents for heart and nervous
 PT system disorders -
 PS PS
 PS Claim 2; Fig 1; 40pp; French.
 CC The present sequence represents a mechanically sensitive potassium
 CC channel protein designated TRAK. The protein is activated by
 CC polyunsaturated fatty acids, particularly arachidonic acid, and by
 CC riluzole. The protein is used to screen for specific modulators which
 CC are useful for treating or preventing diseases of the heart and nervous
 CC systems in humans and animals, e.g. epilepsy, cardiovascular disease
 CC (arrhythmia), neurodegeneration (particularly where associated with
 CC ischemia or anoxia), abnormalities of hormone secretion and muscular
 CC disease. The protein itself may be used to treat these diseases.
 CC Antibodies specific for the protein are used to detect it in tissues,
 CC also as therapeutic inhibitors or activators.
 CC
 XX Sequence 398 AA;
 SQ
 Query Match 21.6%; Score 379.5; DB 20; Length 398;
 Best Local Similarity 34.4%; Pred. No. 1.9e-31;
 Matches 90; Conservative 54; Mismatches 95; Indels 23; Gaps 9;
 Db 18 RSAMCGFLVGLYLVFGAVFSSVELPYEDLLRQELRKIKRRTLEHEHCISEQOLEQ 77
 2 rstllallal-vllylvsgalvfgaleqhegqgkkmhgrdflldhpcvsgksld 60
 QY 78 FLGRVLEASNTGVSVLSNAGSNM-----WDFTSALFPASTVLTSTGCHTVPLSDG 129
 Db 61 fikiivealgg9---anpctswtshsahsawmlgsaffifegltitlygnvlhda 116
 QY 130 GKAFIIVSVIGIPFTLLFLAVVORITVHTRRPVLEH--TWGSKOVVALVHVL 186
 Db 117 grlcltlyalvgiprlfmgldvgdrlgsssl-trgylghaealfkwhvppglvrislsavl 175
 QY 187 LGFVTVSCFFET--PAAVSVLEDMNLFESFPCFISLSTIGLDYVPGEGYNOKFREL 244
 Db 176 --flllgcllflvlpftfysme-swsklealfyivltltvgfdyvgdgltgqn-spa 231
 QY 245 YKIGITCYLLGLLIMLVLET 266
 Db 232 YQPIVWFILGfayfasvltt 253
 RESULT 7
 Y34133
 ID Y34133 standard; Protein; 411 AA.
 XX AC Y34133;
 XX

DT 30-NOV-1999 (first entry)
 XX
 DE Human potassium channel K-Hnov59.
 KW Potassium channel; ataxia; arrhythmia; epilepsy; Bartter's syndrome;
 KW cardiovascular disorder; CNS disorder; renal disorder.
 XX Homo sapiens.
 OS WO9943696-A1.
 XX PD 02-SEP-1999.
 XX PF 22-FEB-1999; 99WO-US03826.
 XX PR 19-JAN-1999; 99US-0116448.
 PR 25-FEB-1998; 98US-0076687.
 PR 07-AUG-1998; 98US-0095836.
 XX
 PA (AYXS-) AYXS PHARM INC.
 XX
 XX Curran ME, Hu P, Miller AP, Rutter M, Wang J;
 WPI: 1999-527591/44.
 DR N-PSDB: 211915.
 XX
 PS Claim 3; Page 104-105; 112pp; English.
 CC This sequence represents the human K-Hnov59 potassium channel.
 CC K-Hnov proteins have a high degree of homology to known potassium
 CC channels and may be alpha subunits, which form the functional channel, or
 CC accessory subunits that act to modulate the channel activity. K-Hnov59 is
 CC a 4 transmembrane domain, 2 pore domain potassium channel. The gene is
 CC located on chromosome 19, determined via PCR chromosomal
 CC localisation using primers Z11939 and Z11940. K-Hnov CDNA
 CC were isolated by extension of expressed sequence tags (ESTs) which were
 CC related but not identical to known human potassium channels. Potential
 CC polymorphisms detected as sequence variants between multiple
 CC independent clones. Potassium channels have critical roles in various
 CC cell types and biochemical pathways. Defective potassium channels are
 CC known to cause four human diseases: episodic ataxia with myokymia;
 CC cardiac arrhythmia (long QT syndrome); epilepsy; and Bartter's syndrome.
 CC As potassium channels are critical components of virtually all cells,
 CC it is likely that abnormal potassium channels are also implicated in
 CC certain renal, cardiovascular and central nervous system (CNS) disorders.
 CC Nucleotides encoding K-Hnov proteins may be used for identifying
 CC homologous or related proteins and the DNA sequences encoding them. They
 CC may be used to produce compositions that modulate the expression and
 CC function of the K-Hnov protein and in studying the biochemical pathways
 CC associated with it. They may also be used for the recombinant production
 CC of K-Hnov protein in fermentation cultures. Additionally, such
 CC nucleotides may be used in gene therapy protocols for the treatment
 CC of diseases associated with abnormal potassium channels.
 CC
 XX Sequence 411 AA;
 SQ
 Query Match 21.6%; Score 379.5; DB 20; Length 411;
 Best Local Similarity 32.5%; Pred. No. 1.9e-31;
 Matches 90; Conservative 62; Mismatches 108; Indels 17; Gaps 9;
 Db 25 FLVGLVLYLVFGAVFSSVELPYEDLLRQELRKIKRRTLEHEHCISEQOLEQFGRVLE 84
 Db 51 flvv--vlylllgatvfkaleqhepsqtrtlvqkqtisqhscevnsclellyqgiva 108
 QY 85 ASNYGVSVLSNAGN-WMDFTSALFPASTVLTSTGCHTVPLSDGKAFIIVSVIG 143
 Db 109 alnagllpntgnslshmdlgssffifagvtvlttfgfnlsprteggkfcfcilgallgip 168

Oy	18	RSAMQFGLVLYGTYLLYLVEGAVFSSVELPYRDXDLROELRKXRRFLREHLELSQQLQ	77
Dd	2	: : : : : : : :	60
Oy	78	FLGRVLEASNTGVSVLNASGNMN--MDFTSALFEASTVLSTTGCHTVPESDGGKAF	134
Dd	61	lkevedalggagadepetnstsnssawdlgaftfgslittlygnvalrldsgrlfc	120
Oy	135	IIVSYTGIPFFLLFTAVNQR-----YHWTRRPVLYFHI RMGFSGKOVAIYHAVIL	187
Dd	121	lfyalvgiprlfglilagvgrlgssalrdngighlea----flfkwhvpelrvlsaml-	174
Oy	188	GFWNVSCFEFI--PAAVSVLEDMDNFLESFECFCISITIGLGIDGYVGEGNQFRRLX	245
Dd	175	-lll-gclllfvllfpfcitycme-dwskleaylivivltltvgyfygvyagadprq- spay	231
Oy	246	KIGTCYLLGLLIAMLVLET	266
		: : : :	
		232 gplvwfwlllqlayfasyltt	252
RESULT	10		
Y94426	ID	Y94426 standard; Protein; 393 AA.	
XX	AC	Y94426;	
XX	DT	04-AUG-2000 (first entry)	
XX	DE	Human h-TRAAK polypeptide #2.	
XX	RW	Human; h-TRAAK; potassium channel polypeptide;	
XX	KM	2P domain potassium channel; neurodegenerative disease; stroke;	
XX	KM	psychiatric disorder; neurological disorder; Gene therapy.	
XX	OS	Homo sapiens.	
XX	PN	WO200026253-A1.	
XX	PD	11-MAY-2000.	
XX	PE	03-NOV-1999; 99MO-GB03634.	
XX	PR	03-NOV-1998; 98GB-0024048.	
XX	PR	07-OCT-1999; 99GB-0023668.	
XX	(SMIK)	SMITHKLIN BEECHAM PLC.	
P1	Chapman CG,	Duckworth DM;	
XX	MP:	2000-365583/31.	
DR	N-PDB;	A27106.	
TF	Novel isolated h-TRAAK polypeptides belonging to the potassium channel		
PT	family of polypeptides, useful for the diagnosis and treatment of		
PT	h-TRAAK related disorders,e.g. depression and schizophrenia -		
PS	Claim 12; Pages 21 and 22; 35pp; English.		
XX	Functional genomics was used to identify h-TRAAK polypeptides and		
CC	h-TRAAK polynucleotides from human tissue samples. h-TRAAK		
CC	polypeptides have homology to the 2P domain potassium channel family of		
CC	polypeptides. The h-TRAAK polypeptides and polynucleotides may		
CC	be used in diagnostic assays for conditions related to h-TRAAK		
CC	imbalance and for identifying agonists and antagonists of h-TRAAK		
CC	polypeptides. The h-TRAAK polypeptides and polynucleotides may also		
CC	be useful for treatment and prevention (e.g. as vaccines) of certain		
CC	diseases, such as pain, psychiatric disorders including depression and		
CC	schizophrenia, neurodegenerative disease including Alzheimer's, stroke		
CC	and head trauma and neurological disorders including migraine and		
CC	epilepsy. The present sequence is human h-TRAAK protein #2.		
XX	Sequence	393 AA;	

Query Match Similarity 21.6%: Score 378; DB 21: Length 393;
Best Local Similarity 33.7%: Pred. No. 2.6e-31;
Matches 88; Conservative 53; Mismatches 98; Indels 22; Gaps

QY 18 RSAMGFGVLVGLVLYLVFGAVVSSVLPYEDLLRQELRKRRFLREHECLSEOLEO 77
Db 2 rstllallal-vlllylsgvalfralceqhpqeqqrgelgvrkrlflrahpovsdqelgl 60
QY 78 FLGRVLEASNTGVSVLSNAGSGMN---WDFISALFFASTVLTSTGYGHTVPLSDGGRKFC 134
Db 61 lkeavedalgggpadpelnstsnshsawdlgsafftsglltlltlygynvalrtdaqrlfc 120
QY 135 IIVSVIGAPFTLLFTFAVORI-----IYVTRRPVLVFIHRGFSKOVAIVHAIVLL 187
Db 121 lfyalvaylpflgillagygdrllgslrlnglhea-----lflkwhypelavysaml- 174
QY 188 GFVTVSCFFET--PAAVESVLEDDWNLFESEFPCFISLSTIGLADPYDGEQYNOKFRELY 245
Db 175 -lllglcillflvlpftfcyme-dwsklealyfvalvltltvgfygdyagadprqd-spay 231
QY 246 KIGITCVLLLGIANLVYLET 266
Db 232 qplvwfwillglayfasvltt 252

RESULT 11
Y30648
ID Y30648 standard; Protein; 370 AA.
AC Y30648;
XX
DT 18-NOV-1999 (first entry)
XX
DE A mechanically sensitive potassium channel protein TREK-1.
XX
KW Mechanically sensitive potassium channel protein; TREK-1;
KM polyunsaturated fatty acid; arachidonic acid; riluzole; heart disease;
RV nervous system disease; epilepsy; cardiovascular disease; arrhythmia;
KM neurodegeneration; ischemia; anoxia; hormone secretion abnormality;
muscular disease.
XX
XX Mus sp.
OS
PN MO9945108-A2.
PD 10-SEP-1999.
XX
PF 23-FEB-1999; 99WO-FR00404.
XX
PR 05-MAR-1998; 98FR-0002725.
XX
PA (CNRS) CNRS CEMT NAT RECH SCT.
PI Honore E, Fink M, Lazdunski M, Lesage F, Duprat F;
XX
XX WPI; 1999-551038/46.
DR N-PSDB; Z10607.
XX
PT New mechanically sensitive potassium channel, used to screen for
PT specific modulators; potential therapeutic agents for heart and nervous
PT system disorders -
XX
XX Claim 3; Page 23-25; 40pp; French.
XX
XX The present sequence represents a mechanically sensitive potassium
CC channel protein designated TREK-1. The protein is activated by
CC polyunsaturated fatty acids, particularly arachidonic acid, and by
CC riluzole. The protein is used to screen for specific modulators which
CC are useful for treating or preventing diseases of the heart and nervous
CC systems in humans and animals, e.g. epilepsy, cardiovascular disease
CC (arrhythmia), neurodegeneration (particularly where associated with

PI Kato S, Kimura T;
 XX WPI: 2000-182694/16.
 DR
 XX
 PT Novel human proteins having hydrophobic domains useful for treating
 PT osteoporosis, Alzheimer's disease, Parkinson's disease, asthma,
 PT multiple sclerosis, rheumatoid arthritis, cancer, anaemia, and stroke -
 XX
 PS Claim 1; Page 245-247; 351pp; English.

XX This sequence represents a human protein of the invention, which has
 CC hydrophobic domains. The DNA sequences can be used as a probe or as a
 CC genetic marker. The protein can also be used as a marker, and to identify
 CC potential genetic disorders. The DNA and protein can also be used as
 CC nutritional sources or supplements. The protein exhibits cytokine, cell
 CC proliferation, cell differentiation activities and induces production of
 CC other cytokines in certain cell populations. The protein also exhibits
 CC immune stimulating or immune suppressing activity. It can be used in the
 CC treatment of various immune deficiencies and disorders, and to treat
 CC infectious diseases caused by viral, bacterial, fungal or other
 CC infections. The protein is also used for treating autoimmune disorders
 CC such as multiple sclerosis, systemic lupus erythematosus, and rheumatoid
 CC arthritis. It is also useful in the treatment of allergic reactions and
 CC conditions such as asthma, and in immune suppression after organ
 CC transplantation. The protein is useful in regulation of haematopoiesis
 CC and consequently in the treatment of myeloid or lymphoid cell
 CC deficiencies. It is also used in compositions for tissue growth or
 CC regeneration. The protein is also used in the treatment of osteoporosis
 CC or osteoarthritis and in the treatment of periodontal disease and other
 CC tooth repair processes. The protein is used in the treatment of nervous
 CC system disorders such as Alzheimer's disease, Parkinson's disease, and
 CC Huntington's disease. They are useful for protection or regeneration and
 CC treatment of lung or liver fibrosis, reperfusion injury in various
 CC tissues, and conditions resulting from systemic cytokine damage. They are
 CC also used for promoting or inhibiting tissue differentiation. They are
 CC also used as contraceptives since they exhibit activin or inhibin related
 CC activities and as a fertility inducing therapeutic. They are used for
 CC treating various coagulation disorders and in treatment and prevention of
 CC conditions resulting from coagulation activities e.g. myocardial
 CC infarction or stroke. They also acts as receptors, receptor ligands or
 CC inhibitors or agonists of receptor/ligand interactions. They are used to
 CC treat inflammatory conditions such as septic shock, sepsis, ischaemia
 CC reperfusion injury, arthritis, and nephritis. They can be used to
 CC prevent tumours.
 CC
 XX
 XX
 XX Sequence 499 AA;

Query Match 20.5%: Score 359; DB 21; Length 499;
 Best Local Similarity 30.1%: Pred. No. 3.5e-29;
 Matches 87; Conservative 56; Mismatches 98; Indels 46; Gaps 11;

QY 26 LVILGYLVGVAVFSSVLEPYEDLRLQEDLRRLREHEECLSEQLODFLGRVLEA 85
 DB 8 ltsalifylaigaalfevleephkkaeknytklhlkfeqlgqgklklevsda 67
 QY 86 SNIVSVLSNAN-GNMWMDFTSALFPASTVLTSTGYGHTVPLSDGKACFLITYGIPF 144
 DB 68 aggvavaltgqtlfnmw--pnamlfaatvltltlgygnvapktpagrlfcfyglfgy 125
 QY 145 TLLELTAV-----VORITVHTVRPVP-----LYFIHMGFSQVVAIYHVALLG 188
 DB 126 cltvisalgtkfggrakrlgqfltkryslkagltctvifivw-----VILNIV-- 176
 QY 189 FVTVSCEFFTPAAVSVLEDDWNLSEFPCFISLTIGLDVYVGEQYGNOKFRRLYKIG 248
 DB 177 -----lppfvmwte-gwmyleglysfististgfdvaygnpsanlyalyyf 226
 QY 249 ITCLLLGL--IAMLV--VLETFCELHE-LKKFRKMYKKDKDEDOYH 292
 DB 227 velwilyglawlsifvmkvsmfvevnhalkkrrrr--rkessfessph 272

RESULT 14
 Y87291
 ID Y87291 standard; Protein; 394 AA.
 XX
 AC Y87291;
 XX
 DT 11-MAY-2000 (first entry)
 XX
 DE Human signal peptide containing protein HSP68 SRQ ID NO:68.
 XX
 KW Human: signal peptide-containing protein; HSP; diagnosis; cancer;
 KW inflammation; cardiovascular disease; anticancer; anti-inflammatory;
 KW antimicrobial; neurotropic; neuroprotective; cardiovascular; hepatotropic;
 KW antiaesthetic; gene therapy; cell proliferation; neurological disorder;
 KW reproductive disorder; developmental disorder; arteriosclerosis;
 KW cirrhosis; psoriasis; acquired immune deficiency syndrome; anaemia;
 KW asthma; Crohn's disease; infection; Alzheimer's disease; schizophrenia;
 KW Parkinson's disease; Huntington's diseases; ovulatory defect;
 KW muscular dystrophy.
 XX
 OS Homo sapiens.
 XX
 PN MO200000610-A2.
 XX
 PD 06-JAN-2000.
 XX
 PF 25-JUN-1999; 99WO-US14484.
 XX
 PR 26-JUN-1998; 98US-0090762.
 PR 31-JUL-1998; 98US-0094983.
 PR 01-OCT-1998; 98US-0102686.
 PR 11-DEC-1998; 98US-0112129.
 XX
 PA (INCYT-) INCYTE PHARM INC.
 XX
 PI Lal P, Tang YT, Gorgone GA, Corley NC, Guegler KJ, Baughn MR;
 PI Akerblom IE, Au-Young J, Yue H, Patterson C, Reddy R, Hillman JL;
 PI Bandman O;
 DR WPI: 2000-160673/14.
 DR N-PSDB; 298176.
 XX
 PT New human signal peptide-containing proteins useful in treatment,
 PT prevention and diagnosis of e.g. cancer, inflammation and
 PT cardiovascular disease -
 PT
 XX
 PS Claim 1; Page 207-208; 327pp; English.
 XX
 CC 298109 to 298242 encode Y87224 to Y87357 which represent the human
 CC signal peptide-containing proteins HSP68-1 to HSP68-134. HSPs have
 CC anticancer, anti-inflammatory, antimicrobial, neurotropic, hepatotropic,
 CC neuroprotective, cardiovascular and antiaesthetic activities, and can
 CC be used in gene therapy. HSPs can be used to treat or prevent disorders
 CC associated with decreased activity or function of HSP. Antagonists of
 CC HSP are used to treat or prevent disorders associated with increased
 CC activity or function of HSP. Such diseases include cell proliferation
 CC (including cancer), inflammation, cardiovascular, neurological,
 CC reproductive or developmental disorders, (e.g. arteriosclerosis,
 CC cirrhosis, psoriasis, acquired immune deficiency syndrome, anaemia,
 CC asthma, Crohn's disease, microbial or other infections, congestive or
 CC ischaemic heart disease, Alzheimer's, Parkinson's or Huntington's
 CC diseases, schizophrenia, ovulatory defects, muscular dystrophy). HSP
 CC nucleic acids can be used for the recombinant production of HSP, for
 CC detecting HSP in standard hybridisation and amplification assays (for
 CC diagnosis and monitoring), in gene therapy, as antisense, triplex-forming
 CC or ribozyme therapeutics, for detecting related sequences or genetic
 CC variations, and for chromosomal mapping. HSP are also used to raise
 CC specific antibodies (Ab) and to screen for agonists and antagonists
 CC (potential therapeutic agents). Ab are used to diagnose, or monitor,
 CC HSP-related diseases (in usual immunoassays), as therapeutic
 CC antagonists, in competitive drug screens, and for purification of HSP
 CC from natural sources.
 XX

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